



EUROPEAN MEDICAL DEVICE REGULATION

Declaration of Conformity

As Legal Manufacturer, we

3M Deutschland GmbH
Health Care Business
Carl-Schurz-Str. 1
41453 Neuss
Germany

hereby declare under our sole responsibility that the following CE marked device(s)

Trade Name	Synthetic Cast Padding
Intended Purpose	The Synthetic Cast Padding is intended for use as padding under standard casting applications
Reference	MW02, MW03, MW04, MW06, MW03-1, MW04-1
Basic UDI-DI	06082232761010000000023CP

are classified per rule 1 Annex VIII of the Medical Device Regulation (EU) 2017/745, as Class I non-sterile devices in accordance with all applicable provisions of the REGULATION (EU) 2017/745 OF THE EUROPEAN PARLIAMENT AND OF THE COUNCIL concerning medical devices.

Margaret Bessenbach
Manager Regulatory Affairs and Quality
Health Care Business EMEA
3M Deutschland GmbH

April 8, 2020

Date

3M is a trademark of 3M.